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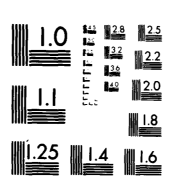
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Photoassisted Water-Gas Shift Reaction over Platinized TiO<sub>2</sub> Catalysts\*

by

S. Sato and J. M. White

Prepared for Publication

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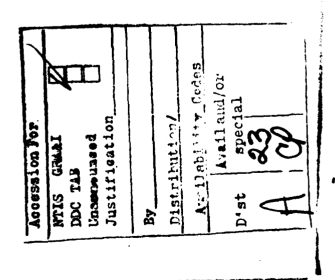
# Photoassisted Water-Gas Shift Reaction over Platinized TiO<sub>2</sub> Catalysts

S. Sato and J. M. White Department of Chemistry University of Texas Austin, Texas 78712

### Abstract

The reaction of gas phase water with CO (water-gas shift reaction) over platinized, powdered  $TiO_2$  is found to be catalytic under UV illumination. The reaction kinetics have been studied at temperatures from 0 to  $60^{\circ}$ C. At  $25^{\circ}$ C, the reaction is zero order both in CO and  $H_2O$  when  $P_2O$  > 0.3 torr and  $P_{H_2O}$  > 5 torr and the activation energy is 7.50 kcal/mole. The wavelength dependence of the reaction rate shows a cut-off near, but slightly below, the band gap of  $TiO_2$ . The quantum efficiency of the reaction is about 0.5% at  $25^{\circ}$ C. A mechanism is proposed which involves the photodecomposition of  $H_2O$  over platinized  $TiO_2$ .

\*Supported in part by the Office of Naval Research.



### I. Introduction.

Heterogeneous photocatalysis is a relatively new branch of catalysis and in almost all cases semiconductors are used because band-gap photon absorption leads to spatially separated electrons and holes which can be used in oxidation-reduction reactions [1]. Metallized semiconductors have recently appeared which are more effective in some cases than semiconductors alone. For example, Bulatov and Khidekel [2] have reported the photodecomposition of acidified water using platinized TiO<sub>2</sub> (Pt/TiO<sub>2</sub>). Kraeutler and Bard [3] have reported the selective decomposition of liquid acetic acid to methane over illuminated Pt/TiO<sub>2</sub>. In the gas phase, Hemminger et al. [4] have claimed the photocatalytic production of methane from CO<sub>2</sub> and gaseous H<sub>2</sub>O using a SrTiO<sub>3</sub> crystal contacted with a Pt foil.

We have recently begun to investigate the photocatalytic properties of TiO<sub>2</sub> and Pt/TiO<sub>2</sub>, and have found that EV-illiminated Pt/TiO<sub>2</sub>, but not TiO<sub>2</sub> alone, catalytically decomposes liquid H<sub>2</sub>O [5]. Platinized TiO<sub>2</sub> also photocatalyzes the reactions of gaseous H<sub>2</sub>O with hydrocarbons [6], active carbon [6] and lignite [7] to produce H<sub>2</sub> and CO<sub>2</sub>, all of which are thermodynamically unfavorable We report here another photoassisted reaction over Pt/TiO<sub>2</sub>, the reaction of gaseous H<sub>2</sub>O with CO (water-gas shift reaction). Although this reaction is not thermodynamically uphill (20° = -6.8 Kcal/mole) over illuminated Pt/TiO<sub>2</sub>, we believe it does involve the photodecomposition of water. Thus the present study, in addition to having intrinsic interest, serves to promote to cur understanding of thermodynamically unfavorable processes over illuminated Pt/TiO<sub>2</sub>.

The purpose of this article is to demonstrate that the photoassisted water-gas shift reaction can be catalytically accomplished over platinized titania at room temperature and that a mechanism which is compatible with the data involves the photodecomposition of water. It is not possible to establish clearly many of the molecular level details of this reaction. This awaits further experimentation involving the combination of optical and electron spectroscopic techniques.

 $TiO_2$  (anatase) was obtained from MCB, doped in flowing  $H_2$  for 6 hr at  $700^{\circ}$ C, cooled in  $H_2$  and stored in a sample vial. Hydrogen was used after passage through a molecular sieve trap at liquid  $N_2$  temperature. Platinized  $TiO_2$  (~2 wtZ Pt) was prepared by the photolysis of hexachloroplatinate dissolved in a  $NaCO_3$ -acetic acid buffer solution [8]. After photolysis the sample was washed with distilled water until  $Cl^-$  could not be detected and then was dried in a desiccator. The BET surface area of  $Pt/TiO_2$  was about  $11 m^2/g$ . Oxygen-18  $H_2O$  (99.37) was obtained from Prochem. Carbon-13 CO (90%) [9] was used after passing through a liquid- $N_2$  trap.

The apparatus consisted of a vacuum system (base pressure 1 x 10<sup>-7</sup> torr), a gas handling system, a reaction system (180 ml volume) and a gas analysis system, most of which were made of Pyrex glass. The reaction system was equipped with a circulation pump, a quartz reaction cell and a storage tube of distilled water which was outgassed several times at dry ice temperature.

The catalyst (0.25 g) was spread uniformly on the flat bottom of the reaction cell and outgassed for 3 hrs at  $200^{\circ}$ C. When studying the water-gas shift reaction, CO was introduced into the reaction system and circulated through the water storage tube to supply water vapor. When low  $\text{H}_2\text{O}$  pressures were needed, the water storage tube was cooled to a temperature at which the desired  $\text{H}_2\text{O}$  vapor pressure was obtained. During reaction the tube was isolated from the reaction system. After

setting the reaction cell temperature with a water bath, the catalyst was illuminated with a 200 W high pressure Hg lamp that was filtered through a quartz cell filled with NiSO<sub>4</sub> solution to remove heat. The gas mixture was sampled at various times, and after passage through a cold trap at about -110°C to remove H<sub>2</sub>O, was analyzed by a mass spectrometer (CEC 21-614). The sensitivity of the mass spectrometer for each gas was calibrated using a gas mixture of known composition.

### 3. Results.

### 3.1 Reaction of gas phase H, 0 with doped TiO,

The hydrogen-doped TiO<sub>2</sub> (without Pt) used in the present experiments reacts with gaseous  $\rm H_2O$  to produce  $\rm H_2$  under UV illumination at room temperature. This reaction was accompanied by the production of a small and variable amount of  $\rm CO_2$  which we believe arises from the photodecomposition of adsorbed carbonate. The rate of  $\rm H_2$  formation dropped to zero after a 2 hr.-irradiation and the maximum amount of  $\rm H_2$  formed was about 3 x  $\rm 10^{-7}$  mole. This reaction was not affected by the presence of CO (i.e. the water-gas shift reaction did not take place) but was completely retarded by the addition of 3 x  $\rm 10^{-2}$  torr of  $\rm O_2$ . Hydrogen was also produced when the TiO<sub>2</sub> was heated in gaseous  $\rm H_2O$  at temperatures above  $\rm 200^{\circ}\rm G$ . There results suggest that the  $\rm H_2$  formation arises from the activated and non-catalytic reaction of  $\rm H_2O$  with a strongly reduced form of TiO<sub>2</sub>, such as Ti<sub>2</sub>O<sub>3</sub> or TiO [10].

### 3.2 Reaction of H20 with Pt/TiQ2

When  $Pt/TiO_2$  was illuminated in the presence of gaseous  $H_2O$ ,  $H_2$  and  $CO_2$  were formed with concomitant formation of a small amount of  $CH_4$ . As shown in Fig. 1, the use of  $H_2^{-18}O$  instead of ordinary  $H_2O$  revealed that a part of the oxygen in  $CO_2$  comes from  $H_2O$ . The ratio of  $(H_2 + 2CH_4)$  to  $(C^{18}O_2 + 1/2 C^{16}O^{18}O)$  is nearly 2:1, suggesting that  $H_2O$  is photo-decomposed on  $Pt/TiO_2$  and that the oxygen formed reacts with adsorbed carbon and CO. These carbon-containing species are probably present as the result of acetic acid decomposition during the preparation of  $Pt/TiO_2$ . The stoichiometric reaction of  $H_2O$  with doped  $TiO_2$ , described in section 3-1, should not contribute to these results since it will have gone to completion during the catalyst preparation. Oxygen atom exchange between  $H_2O$  and  $CO_2$  present originally on  $TiO_2$  may contribute a small amount to the observed  $C^{16}O^{18}O$ . Hydrogenation of adsorbed carbon will account for the formation of  $CH_2$ .

From Fig. 1 the maximum yield of  $\rm H_2$  is certainly larger than 0.09 torr (8.7 x  $10^{-7}$  mole). This is a factor of three larger than the maximum  $\rm H_2$  yield found for  $\rm TiO_2$  and suggests that the addition of Pt increases the activity for  $\rm H_2$  production perhaps through a reaction pathway involving the decomposition of water.

If the photodecomposition of  $\rm H_2O$  is indeed involved,  $\rm H_2$  should be continuously formed by the addition of some material, such as  $\rm CO$ , which has a strong affinity for oxygen and forms a nonreactive product, such as  $\rm CO_2$ . On this basis we tested for, and found, catalytic activity for the water-gas shift reaction on  $\rm Pt/TiO_2$ .

### 3.3 Water-Gas Shift Reaction.

General Features. Figure 2 shows a typical time evolution of the water-gas shift reaction over UV-illuminated Pt/TiO, at 25°C. Carbo 13 CO ( 0.6 torr total, 0.54 torr 13CO) was used in order to discrimina against the CO2 yield that occurs in the absence of CO (g), see Fig. 1. The water pressure was 24 torr. Taking the isotopic purity of CO (90% 1 into account, the stoichiometry of the shift reaction is well  $\Delta p_{\rm H_2}$  . In the dark, the reaction rate satisfied; -Apco = Apco = was very slow at room temperature and, at 60°C, its contribution was only 0.6% of the photo-process. According to Fig. 2, the production of H2 proceeds with no marked loss of activity until the carbon monoxide is consumed. Once the CO (g) is gone, however, the activity drops sharply to zero. In fact the H2 pressure passes through a maximum indicating that some H2 is consumed in the latter stages of the experiment summarized in Fig. 2. In this case the maximum amount of H<sub>2</sub> is 5.8 x  $10^{-6}$  mole (1.3 x  $10^{18}$  molecules  $\pi^{-2}$ of catalyst).

As shown in Fig. 2, a small amount of  $0_2(g)$  appears as the pressure of CO drops below 0.06 torr. This pressure gradually increases at the end of reaction and then abruptly jumps to about  $10^{-2}$  torr when the CO pressure drops below 0.01 torr. No other products were detected in this experiment. The same was true for other reaction temperatures between 0 and  $60^{\circ}$ C.

Reproducibility. The photocatalytic activity was reproducible in a series of runs when the reaction was stopped and the symmetric activity before the CO pressure dropped below about 0.1 torr. If the reaction proceeded to completion, as in Fig. 2, an activity increase

was sometimes observed upon repeating the reaction after evacuation for only a few minutes. Outgassing for extended periods, even at room temperature, restored the activity to a reproducible value. In our experiments reproducible and stable activity was achieved after any experiment by outgassing at 150°C for one hour.

Pressure Dependence at 25°C. Figure 2 shows that the rate of the photoassisted water-gas shift reaction is almost constant until the pressure of CO falls to about 0.3 torr. Over this runge there is a zero order dependence of the rate on CO pressure. Since the pressure of H<sub>2</sub>O (24 torr) is much nigher than the CO pressure (0.6 torr) the effect of its change during the reaction is negligible. The reaction accelerates slightly as the CO pressure drops below 0.3 torr indicating a slightly negative order in p<sub>CO</sub> between 0.3 and 0.1 torr. Near completion the rate slows as the CO pressure drops implying a positive order in p<sub>CO</sub>. These qualitative dependences were confirmed in a series of experiments utilizing initial CO pressures from 0.1 to 0.6 torr.

A slight increase in the initial rate (10 to 20%) was observed when the  $\rm H_2O$  pressure was reduced from 24 torr to about 5 torr while the initial CO pressure was fixed at about 0.6 torr. The reaction is, therefore, almost zero order in  $\rm H_2O$  pressure.

Temperature Dependence. Figure 3 shows the time dependence of the reaction at 0°C. The reaction rate is nearly constant over the entire pressure range observed though a slight decline is observed as the CO pressure drops below 0.25 torr. The consumption

of CO exceeds slightly the formation of  $\rm H_2$  probably due to CO adsorption on the catalyst during the reaction. The  $\rm CO_2$  production rate (not shown) equals the  $\rm H_2$  production rate.

At 60°C (Fig. 4) the time evolution is qualitatively like that at 25°C; an initial constant rate followed by an acceleration at intermediate CO pressures and a retardation at the end of the reaction. However, the CO pressure at which the acceleration begins is higher the in the reaction at 25°C. The decline of the rate is not due to a 1055 of the photocatalytic activity of Pt/TiO because the initial rate was reproduced in the next run. When the initial pressure of CO was reduced to 0.5 torr the rate accelerated earlier that in Fig. 4 and the retardation period became shorter. The reaction course at 40°C showed a pattern intermediate between those at 25°C and 60°C.

Figure 5 shows an Arrhenius plot of the reaction rates for temperatures from 0 to 60°C in the region where the reaction takes place at a constant rate. The activation energy is 7.5 Kcal/mcle. which is about the same as estimated for the photoassisted declars sittion of liquid water (~5 kcal mole<sup>-1</sup>) [5].

Wavelength Dependence. The wavelength dependence was qualitatively measured using three cut-off filters: a commercial UV cut-off filter (415 nm cut-off), a Plexiglass filter (3 mm thick, about 380 nm. cut-off) and a Pyrex glass filter (3 mm thick, about 275 nm cut-off). As seen from Fig. 6, the 415 nm cut-off filter completely retarded the reaction while the Plexiglass and Pyrex filters suppressed the rate to 4 and 67%, respectively. There was no difference in the wavelength dependence at 25 and 60°C.

Formation of 02. The formation of 02 during the water-gas shift reaction is surprising in the sense that it is not favored thermodynamically. The fact that it is formed may help in understanding the machanism of the photoassisted water-gas shift reaction over Pt/TiO2.

typical pressure versus time curves for CO,  ${\rm H_2}$ ,  ${\rm CO_2}$  and  ${\rm O_2}$  are shown in Fig. 7. The amount of  $0_2$  is very small (less than 5 x  $10^{-4}$  torr) and almost constant in the region where the reaction takes place at a constant rate. After the shift reaction accelerates, 0, gradually increases and then maximizes rather abruptly as CO drops below 0.01 torr. At the same time, the H2/CO2 ratio deviates sharply from unity and the excess of  ${\rm H_2}$  over  ${\rm CO_2}$  is about twice the amount  ${\rm O_2}$ formed, suggesting the decomposition of H20. The back reaction occurs readily over Pt and Fig. 7 suggest that the rate of this process increases when the CO pressure drops below  $10^{-3}$  torr. The H<sub>2</sub> consumption rate observed here is much lower than the H<sub>2</sub> oxidation rate on a Pt/TiO<sub>2</sub> serface not exposed to CO [5]. As shown in Fig. 7, the rute becomes a little faster after the light is turned off turning the light source back on produces small amounts of  $\theta_2$  and  $\theta_2$ . This  $\theta_2$  formation pattern was reproducible although the quantitative amounts of 02 produced in each region are sensitive to the activity of Pt/TiO2, the reaction temperature and the initial CO pressure. The maximum amount of (t  $\approx$  25 min. in Fig. 7) is not exactly reproduced but tends to decrease as the reaction temperature increases, as the activity decreases or as the initial CO pressure

is raised.

Preparation Method Dependence. The Pt/TiO<sub>2</sub> prepared from undoped TiO<sub>2</sub> (no H<sub>2</sub>-treatment) showed much lower (~10%) photocatalyti activity than the Pt/H<sub>2</sub>-doped TiO<sub>2</sub> for the water-gas shift and the water decomposition reaction. The activity depends upon the conditions of the H<sub>2</sub>-doping of TiO<sub>2</sub> (temperature, time and a flow rate of H<sub>2</sub>). A strongly reduced TiO<sub>2</sub> leads to higher activity but an optimum condition was not established. As doping proceeds,

TiO<sub>2</sub> changes color from white to light blue, indicating the production of a reduced form of TiO<sub>2</sub>. Preliminary ESCA spectra of the doped and undoped forms revealed no detectable differences.

The Pt loading may also affect the photocatalytic activity but this was not examined in the present study.

### 4. Discussion.

Our experiments show that the photoassisted water-gas shift reaction over Pt/TiO<sub>2</sub> is catalytic when the catalyst is irradiated with band-gap radiation. The reaction may involve photo-induced electronic processes similar to those noted in photoelectrochemical (PEC) cells [11] or photoelectrochemical diodes [12].

At the Pt-TiO, junctions formed by deposition of Pt, the valence and conduction bands will be bent upwards, because charge transfer will occur to equalize the Fermi levels of these materials. The extent to which this occurs for our catalyst cannot be readily determined because the doping level of TiO, is unknown, because the electronic coupling between TiO2 and Pt is not known and because the Pt particles are small and not fully characterizable by bulk parameters. Upon irradiation and excitation of an electron into the conduction band of TiO2, charge separation will compete with hole-electron recombination. Holes will tend to migrate to the Pt-TiO2 interface and the electrons will migrate away from it. Then a photostationary state is reached, the bending of the valence and conduction bands in the semiconductor will be reduced and the electrons and holes can be characterized by quasi-Fermi As a result of these photovoltage levels [11]. effects, we expect (1) that recombination of electrons and holes vill become relatively more important than would be expected on the basis of the "dark" energy level diagram, (2) that holes will be found at both the Pt/TiO, and the TiO,/gas interfaces and (3) that electrons will be captured at the Pt particles. If so, the photocatalyzed processes we have observed here can be described using the diagram (analogous to a shorted electrochemical cell [13]) shown in Fig. 8.

In this mechanism, holes come to the TiO,/gas interface where they react with adsorbed water to form an adsorbed OH radical and a proton. The latter diffuses to the Pt, probably mediated by the physisorbed water that is present, where it is reduced. When CO is present, it reacts with the OH radical to give CO, and a hydrogen atom which recombines with other hydrogen atoms at Pt sites. In the absence of CO, the adsorbed OH radical will either react with a second hole to form H and a chemisorbed oxygen atom or two OH radicals will recombine to form  $H_2O_2$ . Gas phase oxygen would then be formed either by recombination of oxygen atoms or by the rapid decomposition of  $H_2O_2$ . The back reaction,  $H_2(g)$  +  $1/2 \ O_2$  (g)  $\rightarrow$   $H_2O$  (g), occurs readily on Pt [14] and even occurs when liquid water is present [5], but at a slower rate. Consequently, in the absence of CO no measurable O, is evolved. In this case the only way a net amount of H, is evolved is to use surface carbon, present on the catalyst as a result of preparative techniques, to scavenge oxygen and make CO, (See Fig. 1).

The above descriptive mechanism is also applicable to the photolysis of liquid phase water on Pt/TiO2 [5] where the photodissocia-

tion rate is more competitive with the back reaction. There is, however, a problem in view of the fact that the Pt/TiO<sub>2</sub> PEC cell requires an external potential for water decomposition [11,15]. The reason for this requirement is not obvious. There are conflicting views regarding the position of the flat band potential

of TiO<sub>2</sub> with respect to the H<sup>+</sup>/H<sub>2</sub> redox potential. It has been reported as 50 mv positive [11] and 100 mv negative [16]. In the former case, electrons arriving at the Pt electrode of a PEC cell, must pass an energy barrier to reduce protons to hydrogen (this assumes the pressure is 1 atm). In the latter case, the reaction should occur under short circuit conditions. In either case, overvoltages and other voltage drops will contribute to the inefficiency and to the external potential requirements.

Gur results, including those for liquid water [5], show that this reaction does occur without an external potential when the solid photoactive material involves small Pt particles dispersed on powdered TiO<sub>2</sub>. One of the more important differences between our experimental conditions and those in PEC cells is the pressure. The total pressure in our experiments lies between 20 and 25 torn and the hydrogen pressure never exceeds 0.7 torn. The H\*/H<sub>2</sub> redox potential under these conditions will certainly become more positive with respect to the TiO<sub>2</sub> flat band potential, and will allow the reaction to proceed more efficiently. Other differences must also be considered: (1) There will be more surface defects on powdered TiO<sub>2</sub> than on crystalline samples; the attending defect electronic states may be important. (2) The small Pt particles present on our catalyst may have electronic properties different from bulk Pt. (3) Unlike PEC cells, potentially significant [17]

direct chemical bonding between Pt and TiO<sub>2</sub> may be present here. The importance of each of these cannot be established on the basis of our data.

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Another significant factor in the activity of the solid phase is the reduction of the TiO<sub>2</sub> in the preparation procedure. As noted by Wrighton et. al.<sup>15</sup>, this is important in photoelectrochemical cells. One reason is the lower resistivity of the H<sub>2</sub>-doped TiO<sub>2</sub> which enhances photoconductivity. We expect this same change in conductivity to be important in our powdered catalysts. Changes in electronic band structure upon creation of oxygen vacancies are also expected but this can not be examined by kinetic methods. Enhancement of the adsorption of reacting species on the reduced TiO<sub>2</sub> surfaces does not appear to be important.

Focusing now on the water-gas shift reaction, we note that the photodriven dissociation of water is indeed implicated since  $\mathbf{0}_2$  evolution is noted (Figs. 2, 4 and 7). Assuming such a model, the reaction steps probably include:

$$p^{+} + H_{2}O + OH(a) + H^{+}$$

$$p^{+} + OH(a) + O(a) + H^{+}$$

$$CO(g) + CO(a)$$

$$CO (a) + OH(a) + CO_{2} (g) + H(a)$$

$$CO (a) + O(a) + CO_{2} (g)$$

$$CO$$

The intermediates are either denoted in Fig. 8 or described in the text that accompanies it. It is not clear whether Pt participates in steps (3), (4) and (5) since it may be covered with

H(a) during the reaction, as observed in the hydrogen electrode reaction over Pt dispersed on graphite [18].

The time-course of the water-gas shift reaction consists of three regions: (1) region 1 where the pressure of CO is relatively high and the reaction occurs at a constant rate, (2) region 2 where the reaction gradually accelerates with time and (3) region 3 where the depression of the rate and the formation of  $\mathbf{0}_2$  are observed (Figs. 2,4 and 7).

When the CO pressure is high (region 1), the CO coverage on the catalyst surface would be saturated so that the reaction rate is dependent of the CO pressure. We believe this coverage on Ft inhibits  $\mathrm{H}_2$  evolution as observed in the reaction of water with accive carbon [19] and lignite [7] over illuminated Pt/TiO2. The coverage of CO on Pt begins to decrease when the CO pressure is reduced below a certain temperature dependent As a result, the reaction rate accelerates (region 2). When the To pressure is low (region 3), the diffusion of CO through the laver of physisorbed water on TiO, may limit the rate (step 3). As a result the overall reaction rate is suppressed and the coverage of O(a) may increase so that  $O_a$  is produced. Some strongly adsorbed CO would however remain on Pt even after the gaseous CO is consumed and tend to inhibit the reaction of 0, with H2. This adsorbed CO may be slowly removed and allow the back reaction to occur nore rapidly (Fig. 7).

When the reaction temperature is increased, the amount of CO adsorbed on Pt will drop for a given CO pressure. Accordingly, acceleration of the shift reaction starts at higher CO pressures as indicated in Figs. 2-4. The migration of H(a) and O(a) on the catalyst surface, which results in their reaction to form  $\rm H_2O$ , may become important at the end of the reaction at  $\rm 60^{\circ}C$  (Fig. 4).

The reaction order with respect to CO pressure is thus explained in terms of the inhibition by CO (zero and negative order) and the CO diffustion (positive order). The zero order dependence of the rate on  $\rm H_2O$  pressure is reasonable since the catalyst surface would be covered with physisorbed  $\rm H_2O$  even at relatively low  $\rm H_2O$  pressures.

The wavelength dependence of the rate is similar to the wavelength response of the photocurrent in a Pt/W-deped  $TiO_2$  PEC cell studied by Wrighton et.al. [15]. According to their results, a  $TiO_2$  electrode shows a photocurrent threshold near 400 nm and a saturation near 350 nm, while the onset of W-doped  $TiO_2$  is shifted to shorter wavelengths by nearly 50 nm;  $H_2$ -doping and platinizing  $TiO_2$  may have a similar effect.

The quantum efficiency of the present reaction is low (about 0.5% in region 1 at room temperature). This may be improved by changing the level of  $\rm H_2$ -doping and the Pt loading.

### Acknowledgement

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### References and Notes

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### Figure Captions

### Figure 1

Products of the reaction of  ${\rm R_2}^{18}$ 0 with Pt/TiO<sub>2</sub> under UV illumination.  ${\rm P_{H_2}}^{18}_0{\simeq}^24$  torr.

### Figure 2

Time course of the water-gas shift reaction over illuminated Pt/TiO $_2$  at 25°C.  $p_{H_2O} \cong$  24 torr.

### Figure 3

Time course of the photoassisted water-gas shift reaction over Pt/TiO $_2$  at 0°C. Plots of CO $_2$  are omitted as they almost overlap on the plots of H $_2$ .  $P_{H_2O} \cong 5$  torr.

### Figure 4

Time course of the photoassisted water-gas shift reaction over Pt/TiO $_2$  at 60°C. Plots of CO $_2$  almost overlap on those of  $\rm H_2$ .  $\rm P_{H_2}0^{sol}2^4$  torr.

### Figure 5

Arrhenius plot of the rates of photoassisted water-gas shift reaction over Pt/TiO<sub>2</sub>.  $p_{CO} = 0.5 \sim 0.6$  torr,  $p_{H_2O} \simeq 24$  torr (~5 torr to 0°C).

### Figure 6

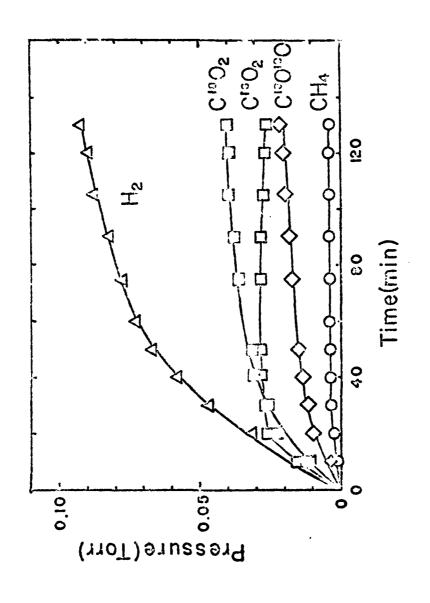
Effects of cut-off filters on the rate of photoassisted water-passistic shift reaction over Pt/TiO<sub>2</sub> at 25 and  $60^{\circ}$ C.  $p_{H_2O} \approx 24$  torr,  $p_{CO} = 0.5 \sim 0.6$  torr.

### Figure 7

Formation of  $0_2$  during the photoassisted water-gas shift reaction over Pt/TiO $_2$  at  $25^{\circ}$ C.  $p_{\rm H_2O}^{\sim}$ 24 torr.

### Figure 8

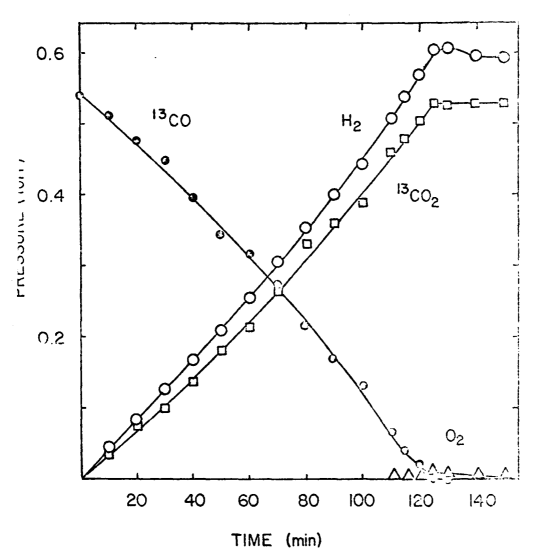
Pictorial Model of the photoinduced redox processes occurring at the  $Pt/TiO_2$  surface.



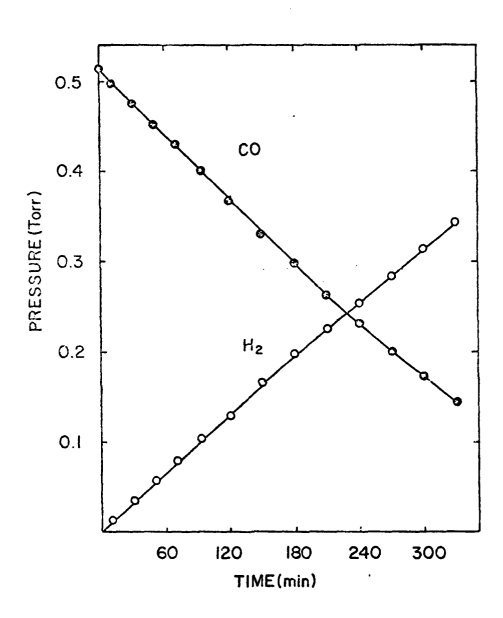
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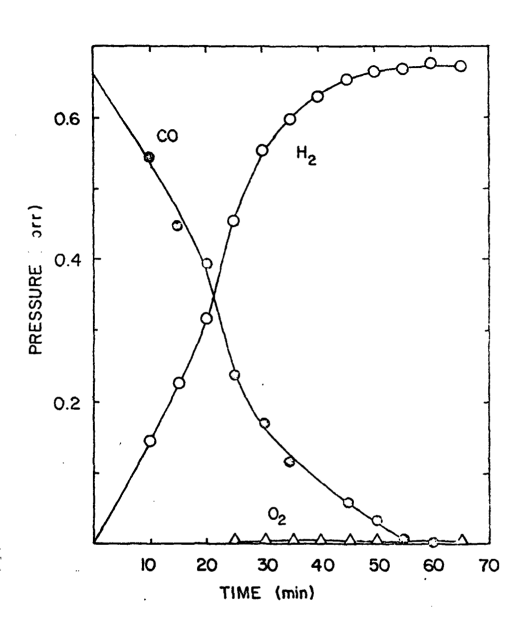




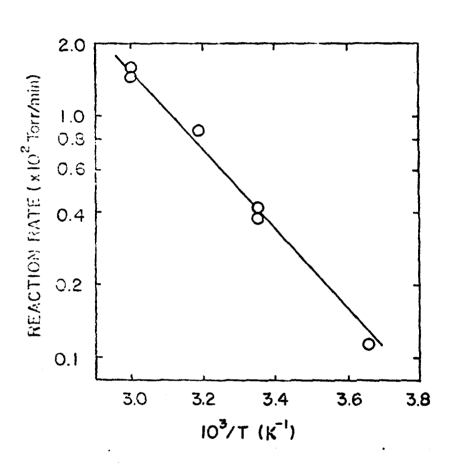
Sato & White, Fig 2.



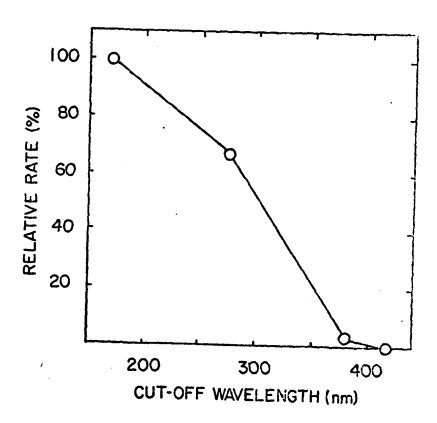
Sato and white. 29.3.



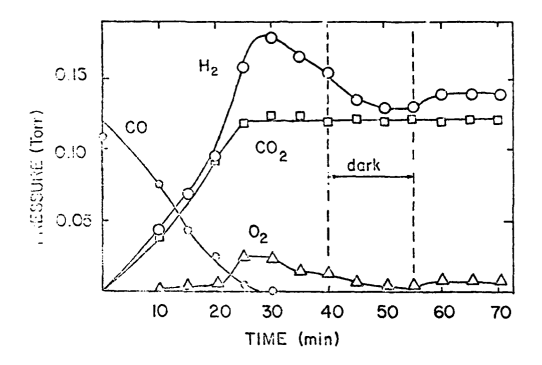
sato & White, Fig. 4.



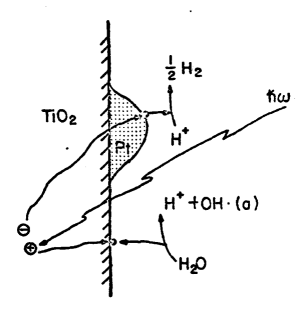
Sato and White, Fig. 5



Sate and white, Fig. 6



Sato & White, Fig. 7



Sato one white , Tiz. 8

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